REMARKS

The Office Action of March 27, 2002 presents the examination of claims 1-10, 14, and 15. No amendments to the application are made and thusly no new matter is inserted into the application.

Request for Interview

If, for any reason, the present invention is not placed into condition for allowance upon entry of this Reply, the Examiner is respectfully requested to contact Kristi L. Rupert, Ph.D. (Reg. No. 45,702) at 703-205-8000 to schedule a personal interview at the Examiner's convenience.

Rejection under 35 U.S.C. § 112, first paragraph

The Examiner maintains the rejection of claim 3 under 35 U.S.C. § 112, first paragraph for allegedly not being described in the specification. Applicants respectfully traverse. Reconsideration of the claims and withdrawal of the instant rejection are respectfully requested.

Claim 3 is directed to the method of claim 2, further comprising determining the presence, position, and type of mutation and categorizing biological aggressiveness and/or metastatic potential of the neoplasia based upon the presence,

position, and type of mutation, wherein said neoplasia is breast cancer, and wherein a mutation in a conserved region II and V of p53 is indicative of poor patient outcome whereas a mutation in a conserved region III and IV is indicative of positive patient outcome.

The Examiner apparently disagrees that one skilled in the art would know that a frameshift or nonsense mutation would be more detrimental than a missense mutation. Specifically, the Examiner writes, "Without guidance or exemplification, one of ordinary skill in the art would not know which type of mutation in conserved region II and V would give rise to a [sic] affect binding or transactivation mutation."

In response to the Examiner's remarks, Applicant submit herewith a journal article, Sjögren et al. "The p53 Gene in Breast Cancer: Prognostic Value of Complementary DNA Sequencing Versus Immunohistochemistry," Journal of the National Cancer Institute, Vol. 88, No. 3/4, 1996 illustrating how mutations which affect DNA binding or transactivation are typically frameshift or nonsense mutations.

In figure 1 of said article, there is a graphic illustration of where mutations are found and the mutation

types. There are essentially four types of mutations that may occur:

- 1. Missense mutations, where a codon is mutated causing a change in the corresponding amino acid.
- 2. Nonsense mutations, where a codon is mutated into a stop codon which terminates protein expression at the mutated codon.
- 3. Deletions which are in-frame, where nucleotides are deleted in multiples of 3 which causes corresponding loss of amino acid(s), or out-of-frame, where nucleotides are deleted in numbers where a premature stop codon is created somewhere downstream of the mutation. In such cases, the expressed protein almost never corresponds to the normal protein.
- 4. Insertions which are in-frame, where nucleotides are inserted in multiples of 3 which causes corresponding addition of amino acid(s), or out-of-frame, where nucleotides are added in numbers where a premature stop codon is created somewhere downstream of the mutation. Again, in such cases, the expressed protein almost never corresponds to the normal protein.

Referring to figure 1 of the paper, it is quite clear that missense mutations are concentrated in the very central, DNA-

binding region of the p53 protein. Other types of mutations are much more widely distributed.

On page 3, lines 9-19 of the specification, it is stated in p53 are missense that approximately 70% of mutations mutations that change the identity of an amino acid and alter the confirmation and stability of p53. Further, on page 7, lines 32-38, it is stated that mutations in p53 that give rise to transcriptional stop signals and a truncated protein prevents p53 from employing its DNA proof-reading role. Finally, on page 8, lines 3-10, it is stated that mutations detrimental to the affect the binding those which DNA patient transactivation, whereas those mutations less harmful for the patient are amino acid changes not greatly affecting structure or function of p53.

Therefore, contrary to the Examiner's remarks, one skilled in the art, given the information disclosed in the specification, would be advised that a frameshift or nonsense mutation would be more detrimental to a cancer patient, whereas a missense mutation would be less detrimental to the cancer patient.

Thus, contrary to the Examiner's assertions, the prognosis of neoplasia based on the "type" of mutation is indeed described

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in the specification so as to reasonably convey to one skilled in the art that the present Inventors had possession of the claimed subject matter at the time of filing.

As these remarks address and overcome the issues of written description raised by the Examiner, Applicants respectfully request withdrawal of the instant rejection.

Rejection under 35 U.S.C. §§ 102, 103

The Examiner maintains the rejection of claim 15 under 35 U.S.C. § 102(e) for allegedly being anticipated by Vogelstein '676 (USP 5,527,676). The Examiner also maintains the rejection of claims 1, 2, 4-10, and 14 under 35 U.S.C. § 103(a) for allegedly being obvious over Vogelstein '676, in view of Elledge et al. and Callahan et al., and further in view of Hedrum et al. Applicants respectfully traverse. Reconsideration of the claims and withdrawal of the instant rejection are respectfully requested.

Claim 15 recites a method for prognostication of the development of neoplasia in a human patient having a neoplasia comprising: a) determining the nucleotide sequence of exons 2-11 of a cancer-related p53 nucleic acid derived from a human neoplastic tissue or body fluid; b) analyzing the entire

nucleotide sequence determined in step a) for the presence of mutations; and c) classifying the neoplasia into different subgroups depending on the presence or absence of a mutation; and d) prognosticating the development of the neoplasia by analyzing the results of step c) only, wherein said results are indicative of patient survival. The Examiner asserts that Vogelstein '676 anticipates claim 15 because "Vogelstein teaches sequencing all of the p53 gene" and "it is inherent that 'part' of the gene that is sequenced encompasses exons 2-11."

method Voqelstein 676 fails to disclose a prognostication of the development of neoplasia in a human Vogelstein '676 merely discloses а method patient. diagnosing a neoplastic tissue of a human (see column 1, lines The difference between the prognostication of the development of neoplasia and the diagnoses of neoplastic tissue are quite different to one skilled in the art, such that the disclosure of one does not destroy novelty of the other.

Typically, the diagnosis of malignant disease on solid tumors is made prior to surgery by conventional means (biopsies, fine needle aspirates) and examined by pathologists looking for typical cellular characteristics to establish diagnosis. In breast cancer at least, this is the sole basis for diagnosis and

initial surgical treatment. After surgical intervention has been made, the primary tumor can be further examined, for instance by: detecting allelic loss of certain defined genes, detecting expression of certain proteins by immunohistochemistry, or by DNA sequencing the entire or parts of genes for mutations.

On the other hand, in order to progosticate a disease i.e., to make a prognosis on how the disease will develop in absence of any additional treatment, as recited in claim 15 in the present application, a number of factors have to be taken into account. In breast cancer, factors that contribute with prognostic information are for instance, nodal status and tumor size. The present invention utilizes the mutational status of the p53 gene in the tumor cells to prognosticate the development of neoplasia. If the prognosis of disease for a given patient is poor, more aggressive treatment is prescribed.

Vogelstein '676 fails to disclose or suggest the prognostication of the development of neoplasia. Further, Vogelstein '676 fails to associate the metastatic potential of the neoplasia based upon the presence, position, and type of mutation. Nor does Vogelstein '676 suggest that certain mutations in p53 are indicative of poor patient outcome.

Instead, Vogelstein '676 merely provides the skilled artisan with methods for assessing p53 in human tumors.

In summary, Vogelstein '676 fails to disclose a method for prognostication of the development of neoplasia in a human patient. As such, Vogelstein '676 fails to anticipate or render obvious the present invention. Further, Elledge et al. and Callahan et al. fail to detect p53 mutations by sequencing exons 2-11 of the gene, whereas Hedrum et al. merely teaches the sequencing of exons 4-9. Thus, absolutely no reference teaches a method for prognostication of the development of neoplasia by sequencing exons 2-11 of p53.

As such, the present invention is not unpatentable over the combination of references cited by the Examiner. Withdrawal of the instant rejection is therefore respectfully requested.

Summary

Overall, the present invention possesses significant patentable features that the cited prior art references do not possess. Furthermore, Applicants submit the instant claims are fully in compliance with 35 U.S.C. § 112, first paragraph. All of the present claims define patentable subject matter such that this application should be placed into condition for allowance.

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Favorable action on the merits of the present application is thereby requested.

Pursuant to the provisions of 37 C.F.R. §§ 1.17 and 1.136(a), the Applicants hereby petition for an extension of three (3) months to September 27, 2002, in which to file a reply to the Office Action. The required fee of \$920.00 is attached to the Notice of Appeal, which is being filed concurrently herewith.

If necessary, the Commissioner is hereby authorized in this, concurrent, and future replies, to charge payment or credit any overpayment to Deposit Account No. 02-2448 for any additional fees required under 37 C.F.R. §§1.16 or 1.17; particularly, extension of time fees.

Respectfully submitted,

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Attachment: Sjögren et al. "The p53 Gene in Breast Cancer: Prognostic Value of Complementary DNA Sequencing Versus Immunohistochemistry," Journal of the National Cancer Institute, Vol. 88, No. 3/4, 1996

a spectrum of community settings as well as men treated at radical prostatectomy warrants further investigations. effectiveness of various follow-up treatment strategies after ties concerning the value of these treatments (14), the low-up cancer treatments after initial surgery and the uncertainselected academic medical centers. Given the wide use of folthe re-treatment rates based on the experience of men treated in treatment decisions, it is important that patients be informed of tional cancer treatment after initial therapy might influence their ing their prostate cancer. Since the likelihood of requiring addibelieve that radical prostatectomy alone is sufficient for managafter radical prostatectomy may have led many patients to remains to be defined (14). Reports of low re-treatment rates cancer. However, its role in the management of prostate cancer Adjuvant therapy is commonly used in the management of

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Notes

¹Editor's nute: SEER is a set of geographically defined, population-based central tumor registries in the United States, operated by local nonprofit organizations under contract to the National Cancer Institute (NCI). Each registry

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annually submits its cases to the NCI on a computer tape. These computer tapes are then edited by the NCI and made available for analysis.

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The p53 Gene in Breast Cancer: Prognostic Value of Complementary DNA Sequencing Versus Immunohistochemistry Versus Immunohistochemistry

Hans Nordgren, Lars Holmberg, Jonas Bergh* Sigrid Sjögren, Mats Inganäs, Torbjörn Norberg, Anders Lindgren,

tions, indicating that IHC failed to detect 33% of the mutabased sequencing method; only 31 (45%) of these mutations analyzed by IHC. The 5-year estimates for RFS, BCCS, and method, and three tumors (1% of the total) could not be sequencing negative (i.e., p53 wild-type), suggesting a 30% tions. Furthermore, 19 of the IHC-positive tumors were tumors (i.e., p53 mutant), 23 exhibited negative IHC reaccoding region. Sixty-four tumors (20% of the total) had were located in evolutionarily conserved portions of the p53 method, and survival curves for different patient subgroups survival (RFS) were estimated by use of the Kaplan-Meier from breast cancer is the considered event), and relapse-free genes expressed in these tumors was screened for mutations cancer specimens. In addition, we determined the prognostic sequencing method and an immunohistochemical (IHC) pose: We compared a complementary DNA (cDNA)-based total) could not be analyzed by the cDNA-based sequencing false-positive frequency with IHC. Four tumors (1.3% of the the presence of mutations. Of the sequencing-positive elevated levels of p53 protein as detected by IHC, suggesting 316 tumors had p53 gene mutations detected by the cDNAvalues are from two-sided tests. Results: Sixty-nine (22%) of were compared by use of the logrank method. All reported P vival (OS), breast cancer-corrected survival (BCCS; death reaction, and DNA sequencing. Probabilities for overall surby combining reverse transcription, the polymerase chain IHC methods. In addition, the entire coding region of p53 nizes both wild-type and mutant forms of p53) and standard use of the mouse monoclonal antibody Pab 1801 (that recogwere evaluated for the presence of mutant p53 protein by used. Methods: Specimens from 316 primary breast tumors value of information obtained when these two methods were method for their abilities to detect p53 mutations in breast alterations has been associated with worse prognosis. Purhuman cancers. In breast cancer, the presence of p53 gene Background: Mutations in the p53 tumor suppressor gene (also known as TP53) have been detected in a wide variety of

in all three categories when compared with those with IHCbreast cancers yielded better prognostic information than method to determine the status of the p53 gene in primary significant. Conclusions: Use of a cDNA-based sequencing negative tumors, but the differences were not statistically Patients with IHC-positive tumors showed reduced survival tive tumors (P = .001, P = .01, and P = .0003, respectively). ing-positive tumors than for patients with sequencing-nega-Natl Cancer Inst 1996;88:173-82] IHC performed with the Pab 1801 monoclonal antibody. [J OS were significantly shorter for patients with p53 sequenc-

influence the induction of apoptosis in malignant cells (6). of the cell cycle (1,4,5). The p53 protein has also been shown to binding properties and an ability to regulate entry into S phase product is a nuclear phosphoprotein. The p53 protein has been cers (3). The p53 gene is located on chromosome 17p, and its detected in a wide range of human tumors, including breast canmany human cancers (1.2). Changes in this gene have been identified as a transcription factor with sequence-specific DNA-TP53) is considered to be a critical step in the development of Alteration of the tumor suppressor gene p53 (also known as

gene have been associated with worse prognosis (7-9) primary, node-negative breast disease, and alterations in the p53 In breast cancer, research has focused on patients with

(IHC). Detection of p53 mutations by IHC is based on the achave used single-stranded conformation polymorphism analysis (SSCP), DNA sequence analysis, or immunohistochemistry Previous studies (8,10-18) evaluating p53 status in cancer

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**See "Notes" section following "References."

approach may provide information more reflective of the funcfore, might be more effective in defining patients with poor biological techniques such as DNA sequencing (21) and, theretional status of the p53 protein than that provided by molecular revealed by IHC, it has been suggested that this methodologic used. Even if the true mutation status of the p53 gene is not two of the antibodies (i.e., antibody 240 and Pab 1801) was antibodies were used and in 45% of the turnors if a cocktail of IHC staining in 18%-35% of the tumors examined if individual comparing five different anti-p53 antibodies revealed positive p53 mutations by IHC has been questioned. A recent study (20) usually detected by IHC. However, the accuracy of detecting (4,19). Wild-type p53 protein has a short half-life and is not changes in the p53 polypeptide that result in increased stability cumulation of mutant p53 protein in cells due to conformational

in relation to adjuvant therapy and prognosis (22). status in the primary tumors of 316 patients with breast cancer study were determined as part of an effort to evaluate p53 gene employing the monoclonal antibody Pab 1801) with that of for p53 gene mutations in breast cancer specimens using IHC In this study, we compared the prognostic value of screening hplete p53 coding region. The cDNA sequences used in this plementary DNA (cDNA)-based sequence analysis of the

Materials and Methods

Study Materials and Patient Population

directly, unfixed and fresh, from the operation theater to the Department of Uppsala County during the study period. All breast cancer samples were sent Sweden. This department contained the only taboratory for histopathology in Department of Pathology, University of Uppsala. Akademiska sjukhuset, December 31, 1989, was used in this study. The tumors were collected at the asive breast cancer in Uppsala County (Sweden) from January 1, 1987, through Tumor material from 316 consecutive patients operated on for primary in-

Lymph node metastases were detected in 97 (31%) of the 316 patients. cars. Tumor sizes ranged from 2 to 130 mm, with a median size of 20 mm, Patient age at diagnosis ranged from 28 to 94 years, with a median age of 63

ears to participate. gram initiated in 1988, which invited all women in Uppsala County aged 40-74 Of the 316 patients, 111 were diagnosed via a mammography screening pro-

node status and p53 data from IHC and cDNA sequencing. clinical data were analyzed together with prospectively recorded data on lymph breast cancer, and 3) death from other causes without signs of relapse. These hree categories: 1) death caused by breast cancer, 2) death with signs of active Patient records were reviewed with regard to relapse information and date of All fatal outcomes were studied, and causes of death were divided into ithout knowledge of the p53 status of the tumors. Details about primary therapy, including radiotherapy, and relapse therapy were also

Therapy and Clinical Follow-up

of high age or the presence of other serious concomitant disease. gissing for 13 patients for whom nodal exploration was not performed because resection or modified radical mastectomy; in both instances, this was combined with axillary lymph node dissection. Axillary lymph node status information is Locoregional therapy. Primary surgical therapy consisted of either sector

derwent sector resection were routinely given radiotherapy, except for those who high age or the presence of concomitant or metastatic disease. Patients who unradiotherapy, except for 19 women who did not receive such therapy because of diameter located in the medial or central area of the breast received locoregional Patients with lymph node-positive disease or tumors larger than 20 mm in

> radiotherapy. [One of these studies has been published (23)]. participated in two randomized studies exploring the efficacy of postoperative

metastatic disease. therapeutic strategies were used for patients with primary inoperable disease or positive disease. Tamoxifen was also given to women with stage II tumors who received intravenous adjuvant polychemotherapy consisting primari ly to all patients with lymph node-positive disease. Premenopausal women ministered. Tamoxifen was given to postmenopausal women with lymph node-When radiotherapy was given concurrently, only cyclophosphamide was adwere lymph node negative as part of a randomized trial. Individualized nine courses of intravenous cyclophosphamide, methotrexate, and 5-Il worouracil. Systemic adjuvant therapy. Systemic adjuvant therapy was offered routineof six to

than the University of Uppsala: they were referred back to the University of through the screening program. sisted of a clinical examination. Blood tests and x-ray procedures were Uppsala (Department of Oncology) on relapse, since this institution has the only least 5 years; after 5 years, they were seen on a yearly basis until 10 years of folformed when indicated. Women aged 40-74 years had mammography checkups clinic for oncology in Uppsala County. The routine follow-up evaluation conlow-up had been completed. A few patients were followed at institutions other seen on a regular outpatient basis at increasing intervals of time each year for at Follow-up. All patients treated for breast cancer in Uppsala County were

Handling of Tumor Material

estrogen- and progesterone-receptor determinations. Corresponding slices from disposable scalpel. Slices were taken from the outer, viable, cellular region of the other half were prepared for histopathologic examination. The part of the one half and frozen in isopentane for later DNA analysis and sequencing and for tumor that was frozen was stored at -70 °C until analysis. Each freshly isolated tumor was divided into two equal portions by use of a

Sequence-Based Analysis of p53 Status

During this procedure, care was taken to avoid thawing of the specimen to microcentrifuge for 10 minutes (at 14 000g), and 350 µL of the aqueous, upper was returned to wet ice for 5 minutes to allow RNA to phase-separate from tisof the tube were then mixed for 10 seconds using a vortex mixer, and the tube Five hundred microliters of RNAzole and 80 µL of a mixture of chloroform and Houston, TX) was placed on wet ice. A section of frozen tumor specimen (5 \times 2 tube containing 300 µL of extraction solution (RNAzole, Cinna Biotec Inc. psala, Sweden). bonate-treated water and 1 μL of RNA guard (25 U; Pharmacia Biotech AB, Up-70% ethanol, dried briefly, and finally dissolved in 50 µL of diethyl pyrocar-The supernatant was discarded, and the pelleted RNA was washed twice with 30 minutes and then subjected to microcentrifugation at 14 000g for 20 minutes phase was recovered and transferred to a new tube that contained 350 µL of sue and other cellular components. Subsequently, the tube was spun in a isoamyl alcohol (in the proportions of 24:1) were added to the tube. The contents tube with a disposable micropestle (Bergman and Beving, Stockholm, Sweden). microcentrifuge tube, and it was pressed and squeezed against the walls of the preparation proved to be necessary. The excised tumor section was placed in the prevent RNA degradation. Such precaution was essential if repeated RNA x 2 mm) to be analyzed was removed with the aid of a disposable scalpel. sopropunol. After brief vortex mixing, the new tube was placed on wet ice for RNA preparation and isolation. A 1.5-mL polypropylene microcentrifuge

at 90 °C for 3 minutes and stored at -20 °C. triphosphatel, and 0.152 A200 U of pd[N], random primers [approximately 2.5 3.6 mM dITP [deoxyinosine triphosphate], 0.9 mM dGTP [deoxyguanosine DTT [dithiothreitol], 3.6 mM dCTP [deoxycytidine triphosphate], 3.6 mM dATP tube containing 10 µL of Moloney murine leukemia virus reverse transcriptase was incubated at 37 °C for 1 hour, and the reaction products were heat-denatured pmol of primers)) to yield a final volume of 75 µL. The cDNA reaction mixture [deoxyadenosine triphosphate], 3.6 mM dTTP [deoxythymidine triphosphate] "cDNA mix" (90 mM Tris-HCl (pH 8.3], 138 mM KCl, 18 mM MgCl₃, 30 mM (200 U, Pharmacia Biotech AB), 2.5 µL of RNA guard (62.5 U), 37.5 µL of 2× tion mixture, 25 µL of a given RNA sample was transferred to a microcentrifuge minutes, followed by chilling on wet ice for 3 minutes. To prepare a cDNA reaccDNA synthesis. The RNA samples were heat denatured at 90 °C for 3

> cling program, followed by incubation at 4 °C minute incubation at 72 °C was performed at the conclusion of the thermocyprofile: 94 °C, 15 seconds; 58 °C, 30 seconds; and 72 °C, 45 seconds. A 5-PCR machine programmed to carry out 38 temperature cycles with the following volumes of 50 µL. The reaction mixtures were incubated in a Perkin Elmer 9600 tive control template was added to specified tubes, yielding total PCR reaction tubes (Perkin-Elmer AB). Five microliters of a given cDNA preparation or nega-(Perkin-Elmer AB), 28 µL of distilled H₂O, and 0.8 µL of Taq polymerase (4 U) the 3' PCR primer (one of them being biotinylated), 1.2 µL of 25 mM MgCl₂ (Perkin-Elmer AB, Sundbyberg, Sweden), 5 pmol each of the 5' PCR primer and (Ampli Taq, Perkin-Elmer AB) were mixed together in individual 0.2-mL PCR Polymerase chain reaction (PCR). Five microliters of 10x PCR II buffer

Design Oligonucleotides (Pharmacia Biotech AB). Four sets of primers were cDNA sequence of p53 messenger RNA. PCR primers were prepared by Custom Primers, PCR and DNA sequencing primers were synthesized based on the

6, 7, 8, and 9 plus parts of exons 5 and 10). Frugment 4: 5'-CGG CGC ACA exons 4 and 7). Fragment 3: 5'-TGG CCC CTC CTC AGC ATC TTA-3' and B-GG-3' (covers the entire sequence of exons 9, 10, and 11 plus a part of exon 8). GAG GAA GAG AAT C-3' and B-5'-CGC ACA CCT ATT GCA AGC AAG 5'-CAA GGC CTC ATT CAG CTC TC-3' (covers the entire sequence of exons GCC AAC CTC-3' (covers the entire sequence of exons 5 and 6 plus parts of B-5'-GTT TCC GTC TGG GCT TCT TGC A-3' and 5'-GGT ACA GTC AGA the entire sequence of exons 2, 3, and 4 plus parts of exons 1 and 5). Frugment 2: TGG ATT GGC-3' and 5'-GCA AAA CAT CTT GTT GAG GGC A-3' (covers used to cover the complete protein coding region of the p53 cDNA. PCR primers (B = biotin-labeled), Fragment 1: B-5'-GAC ACG CTT CCC

DNA sequencing primers (fluorescein-labeled). Fragment 1:F-5'-CAG GGG AGT ACG TGC AAG TCA CAG-3'. Fragment 2:F-5'-GCC AAC CTC CGT-3', Fragment 4:F-5'-GGG GAG CCT CAC CAC GAG CTG-3'. AGG CGG CTC ATA-3'. Fragment 3:F-5'-CGA GTG GAA GGA AAT TTG

an indication that all samples might be contaminated and that the corresponding batch of samples had to be discarded The presence of amplified DNA in negative-control sample tubes was taken as tissue to RNA extraction tubes and not adding RNA to cDNA-reaction mixtures. and the cDNA preparation steps. The negative controls consisted of not adding sequencing, we included negative control samples for both the RNA isolation tion of PCR products that might have originated during the steps preceding DNA Agarose gel electrophoresis and quality control. To control for contamina-

controls were evaluated by subjecting 5-µL aliquots of the relevant PCR the gels as a reference standard bromide. The 100 base-pair ladder (0.2 µg; Pharmacia Biotech AB) was used in products to electrophoresis in 1% agarose gels containing 5 µg/mL ethidium The purity, quality, and quantity of amplified DNA from specimens and from

of plastic combs (solid-phase sequencing combs). coupled Sephurose HP (AutoLoad, Phurmacia Biotech AB) attached to the teeth formed as described by Lagerkvist et al. (24) with the use of streptavidin-Solid-phase DNA sequencing on combs. Sequencing reactions were per-

resulting mixture. PCR products were captured by the comb during an incubation that lasted at least 60 minutes at room temperature. This incubation was inproduct capture (24). terrupted only by occasional raising and dipping of the comb to improve PCR tion of air bubbles), and a solid-phase sequencing comb was immersed in the 2.0 M NaCl). The liquid was carefully mixed by pipetting (avoiding the generataining 80 µL of capture buffer (10 mM Tris-HCl [pH 7.5], 1 mM EDTA, and Forty microliters of a PCR product was transferred to a "4-teeth well" con-

the comb immersed in it, was heated at 55 °C for 5 minutes and then placed at stock solution of fluorescein-labeled sequencing primer were mixed in a third "4-teeth well," followed by insertion of the comb. This annealing mixture, with the comb was washed once with 0.1 M NaOH, once with TE buffer (10 mM 0.1 M NaOH and incubated at room temperature for 5 minutes. Subsequently, room temperature for at least 10 more minutes nealing buffer (AutoRead kit, Pharmacia Biotech AB), and 4 µL of a 1 pmol/mL One hundred four microliters of distilled H₂O, 12 µL of 10x concentrated an-Tris-HCI [pH 7.5] and 1 mM EDTA), and once with ultra-pure distilled H₂O. The comb was then moved to a second "4-teeth well" containing 100 µL of

of T7 DNA polymerase [2 U] [diluted in enzyme dilution buffer, AutoRead kit, concentrated annealing buffer. I µL of extension buffer [AutoRead kit, Phar-Pharmacia Biotech ABI) were dispensed into individual "1-tooth wells" just macia Biotech AB], 4 µL of d/ddNTP mixture, 12 µL of distilled H₂O, and 1 µL Twenty microliters of sequencing reaction mixture (containing 2 µL of 10x

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placed finally on wet ice minutes at 37 °C, and the entire well assembly (with the comb still inserted) was prior to insertion of the comb. The comb was incubated in the wells for 5

apparatus, and ALF electrophoresis was initiated. Biotech AB)-sequencing gel (containing 6% polyacrylamide and 7 M urea) were quencing gel for 10 minutes. The comb was then carefully removed from the gel from the sequencing-reaction wells and inserted into the wells of the ALF-se-STOP solution (AutoRead kit, Pharmacia Biotech AB). The comb was removed pH 8.31), prewarmed to 45 °C, and loaded with 15 µL of 100% formamide rinsed with I x TBE buffer (100 mM Tris, 90 mM bonic acid, and 1 mM EDTA The sample wells of an automated laser fluorescence (ALF; Pharmacia

sequence of a gene with sequences obtained from sample analysis. Evaluator (Pharmacia Biotech AB), version 0.16, which compares the wild-type performed with the aid of a newly developed software program. Sequence Sequencing evaluation and verification. Evaluation of p53 sequences was

further verified by analyzing amplification products from the neighboring cDNA tained within the overlapping portion of two cDNA segments, its existence was quencing the new PCR products. If a mutation was identified in a region con-All mutations were confirmed by reamplifying the relevant cDNAs and se-

Immunohistochemical Analysis of p53 Status

ethanol and distilled water. Pretreatment in a microwave oven at 750 W (three Co., Porthmouth, Philadelphia, PA) were dewaxed in xylene and rehydrated in ecognizes both wild-type and mutant forms of p53, was used at a dilution of mouse monoclonal antibody ct 1801 (12) (Biozac AB, Järfälla, Sweden), which times for 5 minutes each) enhanced p53 antigen accessibility to antibodies. The Paraffin-embedded tumor sections on silane-coated slides (Eire Scientific

tibodies directed against mouse immunoglobulins, avidin-labeled horseradish tions were performed by omitting the primary antibody peroxidase, and DAB as the localization reagents. Negative control IHC reacdiaminobenzidine (DAB) detection kit that includes biotin-labeled secondary antochemistry Instrument (Annex, Helsinki, Finland). We used the manufacturer's Immunostaining was performed in a Ventana ES Automated Immunohis-

considered to be positive. mary IHC analysis, positive p53 staining in any percentage of cancer cells was Positive p53 IHC staining is seen in the nucleus of cancer cells. In our pri-

this multiplied scale at different times without knowledge of clinical outcome from that described by Busch et al. (25). All slides were viewed and judged inyielding a single scale with steps of 1, 2, 3, 4, 6, and 9, where I and 2 were constaining, 2 signified those with positive staining in one third to two thirds of the denoted samples in which less than one third of the tumor cells had positive regard to immunostaining intensity and extent according to graded scales that dependently by two pathologists (A. Lindgren and H. Nordgren) according to and 9 were considered to be high staining. This classification system is derived sidered to be low staining, 3 and 4 were considered to be medium staining, and 6 The results obtained with the two scales were multiplied against each other umor cells, and 3 denoted those with more than two-thirds positive staining tumor cells and 3 represented strongly positive cells. For extent of staining, I ranged from 1 to 3. For intensity of staining, 1 represented weakly positive In a secondary IHC analysis, IHC-positive samples were subclassified with

Statistical Methods

considered to be the event of interest; all other deaths were treated as censoring hazards models. In the multivariate models, age at diagnosis, tumor size sus low variables (cutoff points, 7% for diploid and 12% for aneuploid tumors, sus positive variables (cutoff point, 0.1 fmol/µg DNA) and S phase as high verinto consideration. Hormone-receptor status was dichotomized as negative verrespectively). In breast cancer-corrected survival, death from breast cancer was estrogen- and progesterone-receptor status, and S phase proportion were taken hazards of dying of breast cancer were estimated by use of Cox's proportional of survival curves for different subgroups was evaluated by use of the logrank survivals were estimated by use of the Kaplan-Meier method, and the equality method. All P values are estimated from two-sided statistical tests. Relative Survival probabilities for overall, breast cancer-corrected, and relapse-free

77

causes of death other than breast cancer were excluded. points. Thus, in the breast cancer-corrected survival analysis, all patients with

Clinical Outcome

months, with a maximum follow-up of 87 months. cause of death. The median follow-up in this study was 57 lated causes. For seven patients, we lack information about the cancer, five died with breast cancer present, and 21 died of unre-Of the 316 patients included in the study, 48 died of breast

Mutations Detected and IHC Results

patients could be included in survival analyses. to obtain sequence information; from three others (1% of the study. Since follow-up data were missing for one patient, 308 tumors from 309 patients were available for this comparative total), we lack immunohistochemical data (Table 2). Thus, (Fig. 1). From four tumors (1.3% of the total), we were unable tions were located in evolutionarily conserved regions of p53 insertions (Table 1; Fig. 1). Thirty-one (45%) of the 69 mutaframe deletions, one in-frame insertion, and three out-of-frame premature stop codons), five in-frame deletions, eight out-of-(simple point) mutations, seven nonsense mutations (creating tions were detected in patients whose tumors had metastasized tire protein coding region of the gene. Twenty-nine p53 mutathe 316 patients, p53 mutations were found throughout the encDNA-based sequencing method in tumors from 69 (22%) of Alterations in the p53 gene were detected by means of the lown primary lymph node status. We identified 45 missense illary lymph nodes, 37 mutations were found in node-negalatients, and three mutations were detected in patients with

of the sequence-positive patients (i.e., proven to have p53 muta-(85%) of 13 tumors with deletions (Table 1; Fig. 1). Positive imtions) were negative by IHC, whereas 19 (30%) of the IHCtumors from 64 (20%) of the 316 patients. Twenty-three (33%) tumors with point mutations (Table 1; Fig. 1). munohistochemical reactions were seen in 40 (89%) of 45 with mutations that created premature stop codons and in 11 immunohistochemical reactions were noticed in all six tumors positive patients were sequence negative (Table 2). Negative Positive IHC (suggesting p53 mutation) was demonstrated in

above. There was complete concordance between the two s section) was used to generate the IHC data described rimary method of IHC classification (see "Materials and

> pathologists regarding the assessment of negative and positive immunohistochemical reactions with this method. When the 64 immunohistochemically positive tumors were sub-

versus medium- to high-staining were divergent in only 7.8% of pendent investigators, whereas assessments regarding low-staining grading system, 12.5% of the tumors fell into different low-, pathologists agreed in 75% of the cases. With this more complex classified according to the 6-graded (1-9) scale (i.e., our secondary 53.8% agreement with respect to the proposed class-1 tumors. discrepancy regarding subclasses I and 2, and there was only cases. Within the low-staining group, however, there was a major medium-, and high-staining groups as judged by the two inde-IHC analytical method; see "Materials and Methods" section), the

Comparison of Survival Data

mutation status as determined by IHC and by cDNA-based sepatients), and overall survival (OS; 308 patients). 304 patients), breast cancer-corrected survival (BCCS; 308 quencing. Survival was illustrated as relapse-free survival (RFS; We analyzed the patient data with regard to survival and p53

cording to IHC (Table 3). parameters between p53-positive and p53-negative tumors acnificant differences could be detected in any of the survival statistical significance (P = .2) (Table 3). No statistically sigreduced OS for patients with p53-positive tumors compared with those having p53-negative tumors, but it did not reach IHC and survival. According to IHC, there was a trend of

nificant differences were seen for RFS and BCCS (Table 3). ces in survival were seen between patients with sequencingthose with p53 mutations (P = .0003). Similar statistically sigfor those with positive tumors. The 5-year OS frequency was positive and sequencing-negative tumors, with worse prognosis 18% in the mutation-negative group as opposed to 55% for cDNA sequencing and survival. Highly significant differen-

quencing-positive group (P = .02; Fig. 2, top panel). Differences IHC-positive tumors but negative-sequencing results had died of patients with positive IHC were also statistically significant (62 was 86% for the sequencing-negative group and 56% for the sepanel). The 5-year RFS in this IHC-positive patient population tumors and those with sequencing-negative tumors (Fig. 2, top differences in RFS between those with sequencing-positive patients evaluated, data not shown). None of 19 patients with in BCCS between sequencing-positive and sequencing-negative In 61 women with positive IHC, we saw statistically significant Positive IHC with or without positive cDNA sequencing.

Table 1. Different types of p53 mutations detected by complementary DNA sequencing analysis and corresponding anti-p53 IHC analysis*

	į	D	Deletions		Insertions	D	
	mutations	ln frame	Out of frame	In frame	Out of frame	stop codons	Total
HC +	40	2	0	-	1	0	4
IHC -	s,	w	00	0	_	. 6	23
Unknown IHC	0	0	0	0	_	-	2
Total	45	s	œ	-	w	7	69

IHC = immunohistochemistry.

ARTICLES

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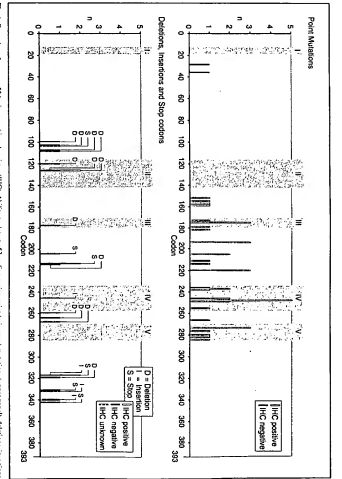


Fig. 1. Detection of mutant p53 by immunohistochemistry (HC). Ability to detect p53 coding region point (missense) mutations (upper panel), deletions, insertions, and premature stop codoms (lower panel) with the monoclonal antibody Pab 1811. Codom positions are outlined below the X axis. The numbers (n) of individual p53 alterations are given on the Y axis. The gray-shaded areas indicate evolutionarily conserved regions 1-V (sere (22) and references contained therein), p53 alterations identified by complementary DNA sequencing but not by HC are indicated by filled black bars, whereas those identified by both methods are indicated by unfilled bars, alterations with unknown HC status are indicated by bars with alternating black and white filling.

top panel). direction for OS, but it was not statistically significant (Fig. 3, the 43 patients with positive-sequencing results had died at 60 breast cancer (BCCS) at 60 months' follow-up, whereas 10 of months of follow-up (P = .03). There was a trend in the same

results. The most marked difference was seen in the OS comparbetween the sequencing-negative and sequencing-positive cases, with worse prognosis for patients with positive sequencing categories, including RFS (243 patients; Fig. 2, bottom panel), statistically significant differences in all three survival In the 246 evaluable patients with negative IHC, there were Negative IHC with or without positive cDNA sequencing.

positive group (P = .002; Fig. 3, bottom panel). quencing-negative group as opposed to 48% in the sequencingison, where the 5-year survival frequency was 78% in the se-

left panel: Fig. 3, left panel) between the IHC-positive and the IHC-negative groups (Fig. 2, were no significant differences in any of the survival categories For the women with sequence-determined p53 mutations, there Positive cDNA sequencing with or without positive IHC.

positive and the IHC-negative groups. There was even a trend nificant differences in survival were observed between the IHC-For the cDNA-based sequencing-negative patients, no sig-Negative cDNA sequencing with or without positive IHC

Table 2. No. of tumors positive and negative for p53 mutation according to IHC- and complementary DNA sequence-based determinations*

			Sequence-base of p	Sequence-based determination of p53 status	
		Mut	Wt	Unknown	Total
	Positive	4	19	_	2
IHC p53 detection	Negative	23	223	w	249
-	Unknown	2	_	0	w
	Total	69	243	4	316

^{*}IHC = immunohistochemistry; Mut = mutation-positive tumors; Wt = wild-type p53.

Yable 3. 5-year survival in relation to p53 mutation status detected by cDNA sequencing and immunohistochemistry (IHC), respectively*

.00L	Positive 54 (66) Negative 71 (238)	p53 status RFS, % (n)	•	
0	5) 69 (66) 38) 86 (242)	t (n) BCCS. 架 (n)	cDNA sequencin	
.0003	55 (66) 78 (242)	OS, % (n)	gra.	5-у
i.	65 (61) 68 (243)	RFS. % (n)		5-y survival
òe	81 (62) 83 (246)	BCCS, % (n)	IHC	
.2	65 (62) 75 (246)	OS, % (n)	**	

*RFS = relapse-free survival; BCCS = breast cancer-corrected survival; OS = overall survival; n = No, of patients; positive = mutation positive by sequencing or positive staining by IHC; negative = wild-type p53 or negative staining by IHC; cDNA = complementary DNA. All P values according to the logrank test.

right panel: Fig. 3, right panel) for better survival in the group with positive IHC reactions (Fig.

Classified as p53 Negative Survival Analysis When IHC Low-Staining Tumors Were

fal between IHC-positive and IHC-negative groups, all 64 our primary IHC analysis of p53 status and survival, which not indicate any statistically significant differences in sur-

tumors with positive IHC staining were considered to be p53 RFS, BCCS, and OS between IHC-positive and IHC-negative tive cases. In comparison with our primary classification, this classes 1-2 (i.e., the "low-staining group" in the more complex from each of the pathologists, considering the IHC-positive subsecondary classification improved all P values for differences in grading system, see "Materials and Methods" section) as negapositive (i.e., mutant). We then reanalyzed the data obtained

Numbers 43 at risk 23

8

Surviving 9,0

0,2

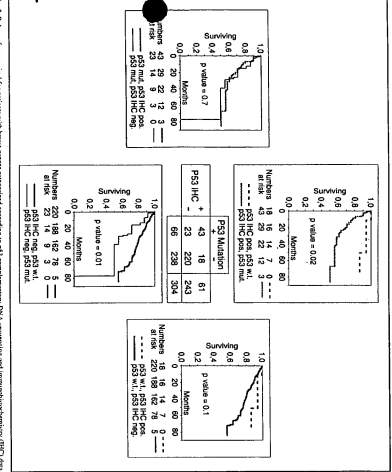


Fig. 2. Relapse-free survival for patients with breast cancer categorized according to p53 complementary DNA sequencing and immunohistochemistry (IHC) data. The four diagrams (top, left, bottom, or right) are based on a 2 × 2 design. Survival curves were generated according to the Kaplan-Meier method; statistical comparisons were made by use of the logrank method. w.t. = wild-type: mut = mutant; neg = negative; pox = positive.

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Fig. 3. Overall survival for patients with breast cancer categorized according to p53 complementary DNA sequencing and immunohistochemistry,(IHC) data. The four diagrams (top, left, bottom, or right) are based on a 2 × 2 design. Survival curves were generated according to the Kaplan-Meier method: statistical comparisons were made by use of the logrank method, w.t. = wild-type: mu = mutant; neg = negative; pos = positive. pathologists, respectively (Table 4). cases was reduced to 36 and 39, as judged by the two p53 mut, p53 IHC pos.
p53 mut, p53 IHC neg. p value = 0.4 34 30 16 18 14 4 8 8 _ ω P53 IHC _ Numbers at risk Numbers 223 207 186 99 at risk 23 18 14 4 --- p53 IHC pos, p53 w.t. p53 IHC pos, p53 mut. Surviving Surviving 0 0 4 6 p53 IHC neg, p53 w.t. p53 IHC neg, p53 mut. <u>0</u>,2 5 9 9 0,2 8,0 6 19 18 16 9 43 34 30 16 p value = 0.002 p value = 0.08 20 40 60 P53 Mutation 8 ಚಿ 66 ည 8 Months †IHC = immunohistochemistry; cDNA = complementary DNA; sequencing *See "Materials and Methods" section for details. 242 Table 4. Overall survival at 5 years in relation to p53 status determined by 223 ඉ 5 cDNA sequencing, IHC (primary analysis*), and IHC modified by 3 ---ප 308 subclassification (secondary analysis*)* Numbers 19 18 16 9 at risk 223 207 186 99 p53 w.t., p53 IHC pos. Surviving O O ဝ 0,2 9,0 5 p value = 0.6 20 40 60 Months 8

this modification in IHC classification the number of positive statistical significance. However, it is important to note that with those from the primary analysis, but none of the values reached the other pathologist's review were also improved relative to statistically significant (Table 4). The P values obtained from data from one of the pathologists, the difference in OS became tumors. The best P values were seen for OS and, according to

Proportional Hazards Models

cDNA sequencing data were higher than those for positive IHC multivariate models (Table 5). The relative hazards for positive monly used prognostic markers. The relative hazards for p53 alclearly included 1.0. data, with confidence intervals indicating an independent effect; terations were of the same magnitude in both the univariate and and IHC was positively or negatively confounded by other comwhether prognostic information generated by cDNA sequencing the confidence intervals for the immunohistochemical data Proportional hazards models were tested to investigate

		5-y overall survival	
	p53	p53 status	
	Positive	Negative	
Method	% (n)	% (n)	P
Sequencing	55 (66)	78 (242)	.0003
IHC	65 (62)	75 (246)	.2
IHC I	52 (36)	76 (272)	.01
IHC 2	59 (39)	75 (269)	.03

= cDNA sequencing; p53 status positive = mutation by sequencing or positive IHC, p53 status regative = wild-type p53 by sequencing or negative IHC. IHC is IHC results by pathologist | (A. Lindgren) after modification with subclasses | and 2 considered loop be negative IHC. IHC 2 = IHC results by pathologist 2 (H. Nordgren) after modification with subclasses | and 2 considered to be negative IHC. All P values according to the logrank test.

Table 5. Results from Cox's proportional hazards models*

Factor	Univariate	Multivariate I	Multivariate
p53 mut versus wt	2.1 (1.1-3.8)	1.9 (1.0-3.7)	t
p53 IHC + versus IHC -	1.2 (0.9-1.5)	l	1.2 (0.9-1.6)
Turnor size	i	1.0(0.99-1.02)	1.0 (0.99-1.0
Node + versus node -	ı	5.0 (2.6-9.9)	5.2 (2.6-10.1
ER + versus ER -	ŀ	1.0 (1.0-1.04)	1.0 (1.0-1.04
PR + versus PR -	!	1.0 (0.93-1.01)	1.0 (0.94-1.0
S phase, high versus low	1	1.5 (0.7-2.9)	1.5 (0.7-3.0)

fect of p53 mutation determined by complementary DNA sequencing; model 2 estimates the relative hazard for IHC data. - = negative; node = axillary lymph nodes; ER = estrogen receptor; PR = progesterone receptor, Estimates of relative hazards with 95% confidence intervals for breast cancer-corrected survival. Multivariate model 1 estimates the ef-*Mut = mutation; wt = wild-type; IHC = immunohistochemistry; + = positive;

changes in exons 5, 6, 7, and 8. contrast with previous investigations of p53 gene alterations in gene were evaluated. Our cDNA sequencing approach stands in of a cDNA-sequencing strategy in which all exons of the p53 accepted (7,16,26-29). The specimens were also analyzed by use whose use on paraffin-embedded tumor samples is now widely with primary breast cancer, was derived from a populationcancer in that the patient population, consisting of 316 women human breast cancer, which have focused primarily on sequence IHC with use of the monoclonal antibody Pab 1801, an antibody based cohort. The breast cancer specimens were examined by his study differs from most other studies of p53 status and

tic information in relation to currently used therapeutic app53. An ideal method should be one that gives the best prognosmethods may indicate that they measure different aspects of lack of concordance between results obtained with these two mutation by cDNA sequencing but were positive by IHC. The detected by cDNA-based sequencing failed to generate positive sible explanations for false-positive and false-negative results results obtained with the two methods, we will now discuss posbased on IHC analyses. In view of the discrepancy between proaches and the best delineation of the patient groups studied IHC reactions with Pab 1801; 19 tumors were negative for with each method. We observed that 23 breast cancers with p53 alterations data clearly demonstrate that the cDNA sequencing method ince most published p53 status determinations have been or in these regards. Our finding should be of impor-

group, where comparison of the sequencing-positive and seresponding IHC-negative group. The suspicion of false-positive not seem to have a significantly worse prognosis than the cortumors had wild-type p53 genes) but positive IHC results did negative results. This was suggested by the observation that the believe that IHC might generate false-positive as well as false-IHC results is supported when considering the IHC-positive patient group with negative cDNA sequencing data (i.e., their quence-based determinations of p53 status gave us reason to The prognostic information generated by IHC- and cDNA se-

> signs of false-negative cases of IHC in the IHC-negative patient those with negative sequencing data. Similarly, in the cDNA sefor the patients with negative sequencing data. There are also quencing-negative patients showed significantly better survival porting our conclusions. between the IHC-positive and the IHC-negative patients, supquencing-positive group, no difference in prognosis was seen patients with positive cDNA sequencing data compared with group, where significantly worse prognosis was observe

six samples with premature stop codons showed negative IHC positive IHC had mutations of these types (P<.001). three tumors with insertions were negative by IHC. Thus, 18 of sible. Our results are consistent with this hypothesis, since all quence of premature stop codons and gross deletions in the p53 ture stop codons. In contrast, only four of 44 samples with quencing but IHC negative had deletions, insertions, and premathe 23 evaluable samples with aberrations detected by cDNA sereactions. Similarly, 11 of 13 tumors with deletions and one of synthesis and render the detection of mutations by IHC imposgene, since such alterations could lead to a cessation of protein False-negative IHC results may be generated as a conse-

mutated protein because of the loss of several important funcof the carboxyl terminus of p53 might reduce the stability of the codons (33), might produce conformational changes in the p53 quencing-positive tumors may be that the genetic alterations bilize the p53 protein sufficiently to be detectable by IHC (21). also possible that certain point mutations may not be able to stathat define the Pab 1801 epitope, supporting these theories. It is codon mutations identified were located downstream of codons detection by IHC impossible. In this study, all deletions and stop cumulation of p53 protein would thus fail to occur, making localization signals, and the oligomerization domain. The actional domains, such as the DNA binding domain, the nuclear polypeptide that interfere with recognition of the epitope for Pab (19,31), as well as deletions, insertions (32), and premature stop Pab 1801 epitope. It has been suggested that missense mutations consequence of premature stop codons should still contain the this region. Furthermore, most of the p53 proteins truncated as a However, few of the alterations that we identified are located in Pab 1801, which is located between amino acids 40 and 65 (30). caused changes in or disappearance of the epitope recognized by 1801. In addition, Ohuc et al. (33) have proposed that truncation Another possible explanation for negative IHC results in se-

data, we observed a nonsignificant trend of better survival for could also represent normal cell cycle fluctuations in p53 sion and mutation status. In some cases, weak immunostaining the existence of a regulatory defect rather than mutations in the in transformed cells (4,19). However, it is possible that the acwild-type p53, which results in the accumulation of p53 protein nals only. For the 243 patients with negative cDNA sequencing type p53 cDNA sequences displayed weakly positive IHC sig-13 of 19 tumors that were positive for p53 by IHC and had wildprotein levels, as indicated in a few reports (36,37). In our study, (34,35) have found discrepancies between p53 protein expresprotein-coding sequence of the gene. Several investigators cumulation of p53 in tumor cells may, in some cases, indicate IHC is that mutant p53 protein exhibits a longer half-life than The theoretical basis for the determination of p53 status by

> chemotherapy, or radiotherapy. which might facilitate apoptosis induced by tamoxifen therapy, be due, in part, to increased amounts of normal p53 protein, tive tumors. One may speculate that the improved survival could those with IHC-positive tumors relative to those with IHC-nega-

as a continuous variable than as a dichotomous variable (8). staining (38). It has also been demonstrated that IHC with Pab biological methods to a greater extent than is low-grade IHC It has been shown that strong immunostaining is associated with the presence of p53 gene alterations detected by molecular 1801 generates better prognostic information when considered Can IHC analysis be refined to increase its resolving power?

data, became statistically significant (OS, IHC positive versus categories. One of the P values, on the basis of one pathologist's nificance of the P values obtained in all three measured survival in which the IHC low-staining group (i.e., subclasses 1 and 2) the method and increasing the risk of false-negative cases. cases was substantially reduced, thus lowering the sensitivity of obtained with the use of sequencing data, the number of positive ulatory system. To achieve a similar level of significance as that and/or more fundamental biological properties of the p53 regnostic specificity of IHC is impaired by technical limitations P values were significant. This result may indicate that the diagtive reactions were considered as p53-positive cases) none of the IHC negative), whereas in our primary analysis (where all posiwere considered to be negative. This approach improved the sig-With these thoughts in mind, we made an additional analysis

to suspect false-positive sequencing data in our study. This conproducts, indicating that the integrity of the tumor isolates was negative cDNA/PCR controls yielded any amplification clusion is further strengthened by the finding that none of the reported by others (39). Given this diversity, we have no reason results may occur as a consequence of contamination of samples based analysis in the same respect. False-positive sequencing and false-positive results, we will now discuss the sequence-14 remaining mutations were located in mutational "hot spots" ferent codons resulting in 55 different mutations. Seven of the tions in 69 cases. Most of these alterations were found in 49 difduring processing. In this study, we identified p53 gene muta-Having analyzed the IHC data with regard to false-negative

p53 coding region with our approach. binding. This risk would be greatest at the extreme ends of the a position disadvantageous for proper primer (cDNA or PCR) might also fail to detect a mutation if the alteration is located in study should have minimized such a risk. cDNA sequencing the wild-type sequence to "drown out" the mutant sequence. malignant cells in relation to normal cells, which could cause tumor samples used for the analysis contained relatively few represent a greater risk. Theoretically, this could happen if the The manner in which the tumor material was isolated in this False-negative sequencing reactions may, on the other hand,

occur with IHC. In terms of using p53 status in clinical decision coding region by use of the present method is taken as the gold prognosis in breast cancer. If complete sequencing of the p53 based analysis of p53 status is superior to IHC in determining standard, false-positive as well as false-negative results can Taken together, our data indicate that direct cDNA sequence-

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making with regard to adjuvant therapy, both false positives and false negatives would pose problems.

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Men and Women: Examination of the Evidence Differences in Lung Cancer Risk Between

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major histologic types were estimated at increasing levels of more precise histologic classification of the cancer type, action of a potential gender difference in relative risk (RR) of but it is likely due to the higher susceptibility to tobacco carmore, this gender difference cannot be explained by difmen at every level of exposure to cigarette smoke. Furtherlung cancer types are consistently higher for women than for epidermoid carcinomas. Adjustments for weight, height, or cinomas and adenocarcinomas than for squamous/ ferences were more pronounced for small-cell/oat cell carcigarette smoking were 1.2-fold to 1.7-fold higher in women contrast, dose-response ORs over cumulative exposure to deeply, and smoked more cigarettes per day than women. In older). Men started smoking earlier, reported inhaling more cer (8.3% for women versus 2.9% for men 55 years old or particularly those with the squamous/epidermoid-type canthat women were more likely to be never-smokers than men, exposure to cigarette smoke. Results: Our results indicated significance was determined by two-sided tests. The ORs for ditional multiple logistic regression analysis, and statistical estimated from adjusted odds ratios (ORs) by use of unconconfounding. The RRs and 95% confidence intervals were and non-Caucasians were excluded from analyses to avoid sex, hospital, and the time of hospital admission. Ex-smokers hospitals from 1981 to 1994 and were pair-matched by age, case and control subjects were admitted to participating and 948 females) with diseases unrelated to smoking. The adenocarcinoma types and 2070 control subjects (1122 males of squamous/epidermoid, small-cell/oat cell, large-cell, and case subjects (1108 males and 781 females) with lung cancer American Health Foundation. It included data from 1889 an ongoing hospital-based, case-control study by the body size. Methods: The present investigation was a part of curate quantitation of smoke exposure, and adjustments for several additional case and control subjects and included lung cancer due to smoking. We added information from up-to-date, more in-depth evaluation of our earlier observatibility to tobacco carcinogens. Purpose: We conducted an biochemical studies suggest gender differences in suscepdecline of smoking among women. Recent epidemiologic and crease in U.S. women exceeds that expected from a slower in U.S. men but is continuing to rise in U.S. women. This in-Background: Lung cancer incidence is gradually leveling off cinogens in women. [J Natl Cancer Inst 1996;88:183-92] ferences in base-line exposure, smoking history, or body size, results confirm our earlier finding that the ORs for major body mass index did not alter the ORs. Conclusions: These than in men for the three major histologic types; these dif-

greater exposure to cigarette tar (1). tinued higher incidence rates in men reflect their longer and cipal cause of lung cancer in both men and women. The con-It is a well-established fact that eigarette smoking is the prin-

the next two to three decades. smoking prevalence among women than among men (1), the exthe leading cause of cancer deaths among U.S. women since States showing that, while lung cancer incidence is leveling of among women are expected to surpass those among men within Consequently, if current trends continue, the lung cancer rates proached and, in fact, may soon surpass that of men posure of women to tobacco carcinogens has gradually ap-1987 (4). At the same time, because of the slower decline in lung cancer mortality since 1950 (3), surpassing breast cancer as women (2). In fact, there has been a 500% increase in female among men, it is continuing to rise at a steady rate among A pattern has evolved during the past decade in the United (2)

is still inconclusive, the potential public health consequences of male and female smoking rates alone. Although the issue of a the rate of decrease in the gap between male-female lung cancer which suggest that, dose for dose, women may be more suscepsuch a phenomenon would be substantial. higher susceptibility to tobacco carcinogens by female smokers than would be expected on the basis of the changing trends in rates observed during the past three decades is more pronounced tible to tobacco carcinogens than men, are of concern. In fact In light of these trends, recent epidemiologic findings (5-13).

on as many as seven different brands of cigarettes smoked. titation of lifelong smoking exposure for each participant based study participants. As a result, we now have more precise quanstarted collecting more detailed smoking histories from the susceptible to tobacco carcinogens than men, was limited to a finding by using more precise histologic subtypes. In 1985, we and Kreyberg II types. It is important to further evaluate this broad histologic classification of lung cancer, i.e., Kreyberg I Our previous work (8), suggesting that women may be more

ducted an in-depth evaluation of the differences in lung cancer posures and more defined lung cancer histologies), we con-(with more detailed and precisely quantitated smoking exadditional data on more case subjects as well as control subjects risk between men and women. By reviewing the results of Spurred by our initial findings (8) and by the availability of

ARTICLES

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